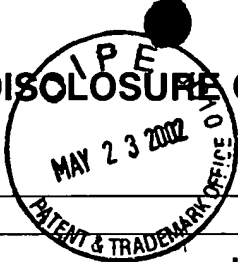


INFORMATION DISCLOSURE CITATION				Docket No.: RLL 03US	Serial No.: 09/998,115
				Applicants: Anand, et al.	
				Filed: 11/30/2001	Group:



U.S. PATENT DOCUMENTS

EXAMINER INITIAL		DOCUMENT NUMBER	DATE	NAME	CLASS	SUBCLASS	FILING DATE IF APPROPRIATE
FB	A1	4,479,954	10/30/1984	Hirose et al.	424	251	TECH CENTER 1600/2900 MAY 24 2002
FB	A2	4,524,206	6/18/1985	New et al.	544	230	
FB	A3	4,598,078	7/1/1986	Ishizumi et al.	514	252	
FB	A4	6,083,950	7/4/2000	Anand, et al.	514	252	
FB	A5	6,090,809	7/18/2000	Anand, et al.	514	252	

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FOREIGN PATENT DOCUMENTS

		DOCUMENT NUMBER	DATE	COUNTRY	CLASS	SUBCLASS	TRANSLATION YES NO
FB	B1	WO 98/37893	9/3/1998	PCT			X
FB	B2	WO 98/51298	11/19/1998	PCT			X
FB	B3	JP 59036661	2/28/1984	Japan			X Abstract only
FB	B4	JP 60-204784	10/16/1985	Japan			X Abstract
FB	B5	JP 02-235865	9/18/1990	Japan			X Abstract
FB	B6	EP 0 109 562 A1	5/30/1984	EPO			
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FB	B8	EP 0 711 757 A1	5/15/1996	EPO			
FB	B9	AT 387 773 B	3/10/1989	Austria			X

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FB	C1	HIEBLE et al., "Recent advances in the identification of alpha1- and alpha2-adrenoceptor subtypes: therapeutic implications", <u>Expert Opinion on Investigational Drugs</u> 6, pp. 367-387 (1997)
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*EXAMINER: Initial if reference considered, whether or not citation is in conformance with MPEP 609; Draw line through citation if not in conformance and not considered. Include copy of this form with next communication to applicant.			

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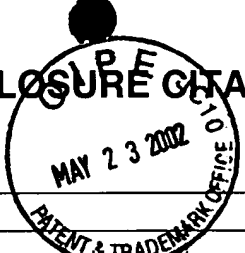
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<i>FB</i>	C3	CROSS, et al., "Substituted trifluoromethyl phenyl piperazines as anoretic agents", <u>Eur. J. Med. Chem. – Chimica Therapeutica</u> <u>12</u> , pp 173-176 (1977)
<i>FB</i>	C4	VAN STEEN, et al., "A Series of N4-Imidoethyl Derivatives of 1-(2,3-Dihydro-1,4-benzodioxin-5-yl)piperazine as 5-HT _{1A} Receptor Ligands: Synthesis and Structure-Affinity Relationships", <u>J. Med. Chem.</u> <u>38</u> , pp. 4303-4308 (1995)
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<i>FB</i>	C6	NEW, et al., "Buspirone Analogues. 2. Structure-Activity Relationships of Aromatic Imide Derivatives", <u>J. Med. Chem.</u> <u>29</u> , pp. 1476-1482 (1986)
<i>FB</i>	C7	KORGAONKAR, et al., "Synthesis of N-[3-(4-Aryl-1-piperazinyl)propyl]-4, 4-bis(4-methoxyphenyl)piperidine-2,6-diones/Tetrahydropthalimides/Camphorimides as Sedatives", <u>J. Indian Chem. Soc.</u> Vol. LX, pp. 874-876 (1983)
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<i>FB</i>	C10	SAMANT & KULKARNI, "Synthesis and Pharmacology of N-(N ⁴ -Aryl-N ¹ -Piperazinylalkyl) Phthalimides : CNS Depressants", <u>J. Indian Chem. Soc.</u> Vol. LVI, pp. 1002-1005 (1979)
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<i>FB</i>	C15	ZAGIDULLIN, "N-(beta-Aminoethyl)piperazine and its derivatives in aminomethylation reactions", <u>Zh. Obshch. Khim.</u> , vol. 16, no. 1, pp. 247-253 (1991)
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